# Activity Outline FDA Drug Topics: Orange Book: Frequently Asked Questions and Answers November 16, 2020 FDA

**Activity Coordinator:** 

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#### **Series Description**

This series of educational webinars is designed to aid physicians, physician assistants, nurses, pharmacists, pharmacy technicians, certified public health professionals, other health care professionals, and students, to provide better patient care by knowing how to find relevant FDA regulatory information that will improve drug safety.

#### **Lecture Description**

This webinar is intended to assist stakeholders in utilizing the Orange Book. It will address some of the most frequently asked questions that the FDA has received from interested parties regarding the Orange Book.

#### References

- Orange Book Questions and Answers Draft Guidance for Industry (May 2020) (https://www.fda.gov/regulatory-information/search-fda-guidance-documents/orange-book-questions-and-answers-guidance-industry)
- Orange Book (www.fda.gov/OrangeBook)
- Orange Book Preface (https://www.fda.gov/drugs/development-approval-process-drugs/orange-book-preface)
- Additional Orange Book References (https://www.fda.gov/drugs/drug-approvals-and-databases/approved-drug-products-therapeutic-equivalen ce-evaluations-orange-book)

# **Series Objectives**

- Explain how to utilize FDA's drug information, medication safety resources, and regulatory guidance, to improve delivery of patient care and optimize outcomes.
- Describe and inform health care providers of recent labeling, policy and regulatory changes which would impact prescribing and medication management to optimize patient care.

# Learning Objectives After completion of this activity, the participant will be able to:

- Utilize the Orange Book to identify therapeutically equivalent products
- Identify the regulatory requirements for submission of patent information to the FDA
- Recognize the statutory requirements for the movement of drug products between the active and discontinued sections of the Orange Book and the timing for publication
- Employ the Orange book website to locate relevant regulatory information

#### **Target Audience**

This activity is intended for physicians, pharmacists, pharmacy technicians, nurses, CPH - Certified Public Health, and physician assistants.

# Agenda

## Lecture 1 November 16, 2020

Time	Topic	Speaker
1:00 - 2:00 PM	Orange Book: Frequently Asked Questions and Answers	Kendra Stewart

#### **Continuing Education Accreditation**



In support of improving patient care, FDA Center for Drug Evaluation and Research is jointly accredited by the Accreditation Council for Continuing Medical Education (ACCME), the Accreditation Council for Pharmacy Education (ACPE), and the American Nurses Credentialing Center (ANCC) to provide continuing education for the healthcare team.



This activity was planned by and for the healthcare team, and learners will receive 1 Interprofessional Continuing Education (IPCE) credit(s) for learning and change.

# **CME**

FDA Center for Drug Evaluation and Research designates this live activity for a maximum of 1.00 *AMA PRA Category 1 Credit(s)*™. Physicians should claim only the credit commensurate with the extent of their participation in the activity.

#### CPE

This knowledge-based activity has been assigned ACPE Universal Activity Number JA0002895-0000-20-006-L04-P, and ACPE Universal Activity Number JA0002895-0000-20-006-L04-T for 1.00 contact hour(s).

#### CNE

FDA Center for Drug Evaluation and Research designates this activity for 1.00 contact hour(s).

#### **AAPA**

This activity is designated for 1.00 AAPA Category 1 CME credits. FDA Center for Drug Evaluation and Research has been authorized by the American Academy of PAs (AAPA) to award AAPA Category 1 CME credit for activities planned in accordance with AAPA CME Criteria. PAs should only claim credit commensurate with the extent of their participation.

#### **CPH**

Up to 1.00 CPH Recertification Credits may be earned at this event.

# Requirements for Receiving CE Credit

Physicians, physician assistants, pharmacists, nurses, pharmacist techs, and those claiming non-physician CME:participants must attest to their attendance and complete the final activity evaluation via the CE Portal (ceportal.fda.gov). For multi-day activities, participants must attest to their attendance and complete the faculty evaluation each day. Final activity evaluations must be completed within two weeks after the activity - no exceptions.

Attention Pharmacists and Pharmacy Techs: Failure to provide your correct NABP AND Date of Birth information, in the required format, may result in the loss of credit for this activity. NABP profile number should be the 6-7 digit profile number assigned by the CPE Monitor and your birth date should be in the MMDD format (e.g. 0721) Do not provide your pharmacy license number. Please click the "My Account" tab and then navigate to "Edit Contact Information" to verify that your information is correct.

## Important Note regarding completion of evaluations and receiving credit

Attendees have 14 days from the last day of the activity to log in, complete the required evaluation(s) and attest to your attendance to claim credit. Physicians, physician assistants, and nurses may then view/print statement of credit. Pharmacists should log into the CPE monitor 10 weeks after the last session of the activity to obtain their CE credit.

## **Disclosure**

# **Faculty**

□ Stewart, Kendra, Supervisor, Orange Book, CDER - nothing to disclose

#### **Planning Committee**

- Burke, Kara, PharmD, Team Leader/Pharmacist, FDA/CDER/OCOMM/DDI nothing to disclose
- □ Cao, Christian, MPAS, PA-C, Safety Evaluator Team Leader, FDA/CDER/OSE/DPV nothing to disclose □ DeFronzo, Kimberly, RPh, MS, MBA, Consumer Safety Officer, FDA/CDER/OCOMM/DDI nothing to disclose
- □ Kapoor, Rama, MD, Medical Officer, FDA nothing to disclose
- Navin, Lesley, RN, MSN, Consumer Safety Officer, FDA/CDER/DDI nothing to disclose
- □ Nguyen-Chu, Thanh Tam, PharmD, Pharmacist, FDA/CDER/OCOMM/DDI nothing to disclose

# **CE Consultation and Accreditation Team**

- □ Lisa Thompson, MSHA, MBA, CE Consultant, FDA/CDER/OEP/DLOD nothing to disclose
- □ Zawalick, Karen, CE Team Leader, FDA/CDER/OEP/DLOD nothing to disclose

# **Registration Fee and Refunds**

Registration is complimentary, therefore refunds are not applicable.